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Intersystem crossing processes in the 2CzPN emitter: a DFT/MRCI study including vibrational spin–orbit interactions†

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Multireference quantum chemical calculations were performed in order to investigate the (reverse) intersystem crossing ((R)ISC) mechanisms of 4,5-di(9*H*-carbazol-9-yl)-phthalonitrile (**2CzPN**). A combination of density functional theory (DFT) and multireference configuration interaction methods (MRCI) was used. The excellent agreement of the computed absorption spectrum with available experimental absorption spectra lends confidence to the chosen computational protocol. Vertically, two triplet excited states (T_1 and T_2) are found below the S_1 state. At the excited state minima, the calculated adiabatic energies locate only the T_1 state below the S_1 state. The enhanced charge transfer (CT) character of the geometrically relaxed excited states causes their mutual (direct) spin–orbit coupling (SOC) interaction to be low. Contributions of vibronic SOC to the (R)ISC probability, evaluated by a Herzberg–Teller-like procedure for a temperature of 300 K, are small but not negligible. For ISC, the $S_1 \rightarrow T_1$ channel is the fastest ($8 \times 10^6 \text{ s}^{-1}$), while the $S_1 \rightarrow T_2$ channel is found to be thermally activated ($9 \times 10^4 \text{ s}^{-1}$) and less efficient when proceeding from the adiabatic S_1 state. Our calculations also reveal, however, a barrierless $S_1 \rightarrow T_2$ ISC pathway near the Franck–Condon region. RISC is found to essentially proceed via the $T_1 \rightarrow S_1$ channel, with a rate constant of ($3 \times 10^4 \text{ s}^{-1}$) if our adiabatic singlet–triplet energy gap in vacuum ($\Delta E_{ST} = 0.12 \text{ eV}$) is employed. Shifting the potentials to match two experimentally reported singlet–triplet energy gaps in toluene ($\Delta E_{ST} = 0.21$ and 0.31 eV , respectively) leads to a drastic reduction of the computed rate constant by up to 4 orders of magnitude. The T_2 state is not expected to play a major role in mediating triplet–singlet transitions in **2CzPN** unless it is directly populated by hot excitons. No indication for a strong vibronic coupling of the T_2 and T_1 potentials is found, which could help overcome the negative exponential dependence of the RISC rate constant on the magnitude of the energy gap.

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1. Introduction

The carbazolyl dicyanobenzene (CDCB) family of compounds has been widely investigated as thermally-activated delayed fluorescence (TADF) emitters in organic light emitting diodes (OLEDs).¹ The devices, built using these prototype compounds have shown high luminescence efficiencies and excellent operational stability.^{2–8}

As first proposed by Uoyama *et al.*,⁹ CDCB emitters were constituted by carbazole (Cz) units as donors (D) and the dicyanobenzene (DCB) moiety acting as an acceptor (A). These systems showed strong intramolecular charge-transfer (ICT)

emission with the luminescence colour varying (from sky blue to orange) with the number and relative position of the Cz and cyano groups in the DCB unit. Of this series, the sky blue emitter **2CzPN** (4,5-di(9*H*-carbazol-9-yl)-phthalonitrile, Fig. 1) had the smallest external quantum efficiency (EQE) of 8.0% in toluene and 2,8-bis(diphenylphosphoryl)-dibenzo[*b,d*]thiophene (PPT)

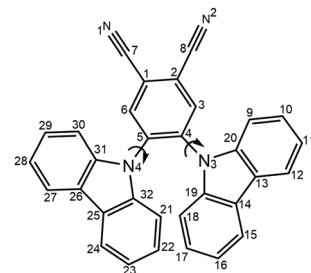


Fig. 1 Chemical structure of 4,5-di(9*H*-carbazol-9-yl)-phthalonitrile (**2CzPN**).

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film, while the green emitter **4CzIPN** showed the highest external quantum efficiency (EQE) in toluene and 4,4'-*N,N*'-dcarbazole-biphenyl (CBP) film (19.3%).⁹ The lower efficiency of **2CzPN** has been attributed to a large roll-off with current increase, which is dominated by strong contributions of singlet-triplet annihilation (STA) and triplet-triplet annihilation (TTA) as shown by Masui *et al.* by using an exciton-quenching model.¹⁰ Later, its efficiency was further improved by including a mixed co-host system, which included electron/hole transporting materials in the device.¹¹ This architecture improved its efficiency to an EQE of 21.8% that is stated to be one of the highest for blue TADF OLEDs.

In TADF, the chromophore exhibits delayed fluorescence (DF) from the lowest-lying (S_1) singlet state, which is thermally populated from an energetically close-by triplet excited state (T_n) by reverse intersystem crossing (RISC). Therefore, with RISC being the rate limiting process in TADF, a small singlet-triplet energy gap (ΔE_{ST}) in the range of the thermal energy in conjunction with an effective spin-orbit coupling (SOC) between target states is crucial for facilitating the up-conversion of the triplet population into emissive singlets.¹² This is particularly critical for **2CzPN** and its derivatives,^{5,6} where the reported ΔE_{ST} values are large compared to those of other TADF emitters.¹ For **2CzPN**, ΔE_{ST} values between 0.21 eV¹³ and 0.31 eV⁷ have been determined in toluene solution yielding RISC rate constants (k_{RISC}) of 6×10^3 s⁻¹ and 1.7×10^5 s⁻¹, respectively. The latter value is remarkably high in view of the sizable singlet-triplet splitting. An exceptionally small ΔE_{ST} value of 0.09 eV was determined in a 1,3-bis(*N*-carbazolyl)benzene (mCP) OLED, yielding a k_{RISC} value of 5.6×10^3 s⁻¹.¹⁰ These observations bring to light interesting optical properties of these emitters that are influencing their k_{RISC} and, consequently, the TADF efficiencies.

The optical and electronic properties of the sky blue **2CzPN** emitter have been extensively studied in solution^{5,11,13} and in a variety of OLED ensembles.^{8,10,11,14-17} In toluene, hexane and dichloromethane (DCM) solutions, **2CzPN** shows similar ultraviolet absorption spectra, which are constituted by three bands: two bands (a maximum and a shoulder) in the range of 300 and 350 nm that were assigned to LE on the Cz moiety and a wide CT band peaking at *ca.* 375 nm.^{11,13,16} The absorption spectrum in tetrahydrofuran (THF) solution is found to be blue-shifted by a few nanometers.⁵ The emission of **2CzPN** shows up at *ca.* 475 nm in toluene and is blue-shifted to 447 nm in hexane,¹⁶ while this peak is red-shifted to 501 nm in diluted THF solution.¹³ Bathochromic shifts of the emission maxima with increasing solvent polarity are typical for CT transitions in liquid solution. For all solvents, the reported rate constant for the radiative decay is similar ($\sim 10^7$ s⁻¹).

Experimental works supported by theoretical calculations have played a significant role in rationalizing mechanistic details about the (R)ISC of TADF emitters. A D-A type of architecture in multi-chromophoric systems leads to a large spatial separation between the frontier molecular orbitals (HOMO and LUMO) and thus promoting a small ΔE_{ST} . Consequently, their lowest-lying excited states are typically of charge

transfer (CT) character. According to the empirical El-Sayed's rules for SOC, the coupling between singlet and triplet states of the same character is forbidden, *e.g.* between ($\pi\pi^*$) CT states. Therefore, vibrationally induced SOC becomes of great importance in mediating efficient (R)ISC for TADF emitters, which in fact has been elegantly demonstrated in experiments and *via* theoretical calculations.¹⁸⁻²⁴ Vibronic coupling can boost the coupling between two ($\pi\pi^*$) CT states by borrowing its intensity from a close-lying state with a local excitation (LE) character. This state can typically bridge the energy gap between the lowest singlet and triplet states as shown by Gibson *et al.*²⁰ and Etherington *et al.*²² for D-A and D-A-D TADF active compounds between phenothiazine as a D and dibenzo thiophene-S-S-dioxide as an A. The influence of multiple excited states and vibrational effects in SOC and their role in TADF has been recently discussed by Marian and Penfold *et al.*^{25,26}

For **2CzPN**, previous studies have demonstrated that conformational freedom modulates the degree of admixture of the LE character in the S_1 and T_1 (CT $\pi\pi^*$) states. These include experimental transient absorption spectroscopy (TAS)¹³ and transient electron-spin resonance²⁷ as well as several theoretical TDDFT-based approaches²⁸⁻³³. Such studies focus on substitution effects on the electronic and excited state properties of several CDCBs performed mainly on ground state geometries. Other computational works calculate first-order SOC values in order to obtain (R)ISC rate constants *via* semiclassical Marcus theory using the adiabatic T_1 state^{34,35} or involving the singlet-triplet crossing seam³⁶.

In this study, we aim at understanding the underlying excited state decay mechanisms that drive the (R)ISC process in **2CzPN**. To this end, we carried out extensive hybrid density functional theory/multireference interaction (DFT/MRCI) calculations. The DFT/MRCI method^{37,38} has been largely used in our laboratory for getting mechanistic insights into (R)ISC of organic³⁹ and inorganic^{40,41} compounds. Furthermore, we give a detailed overview on the (R)ISC efficiencies and their implications for TADF emission.

2. Computational details

As it is well known, substantial errors are obtained when calculating excitation energies of CT states with time-dependent density functional theory (TDDFT) and standard exchange-correlation (xc) functionals. Several computational studies have evaluated the performance of TDDFT for a wide range of TADF emitters (including **2CzPN**) correlating (vertical) singlet and triplet transition energies with experimental absorption energies. This, with the purpose of finding an optimal percentage of Hartree-Fock (HF) exchange contribution to the xc functional that allows the evaluation of transition energies for CT states and singlet-triplet energy gaps in a more precise manner.⁴²⁻⁴⁴ This empirical calibration seems to work very well for states with strong CT character but has limited applicability for mixed CT-LE states and in cases where the LE contributions to a desired state vary with geometry distortions. A more promising ansatz toward a balanced



description of CT and LE states in donor–acceptor compounds in TDDFT calculations is optimal tuning where the ω parameter of a range-separated functional is adjusted such that the negative HOMO energy equals the molecular vertical ionisation.^{45,46}

In this context, we performed an evaluation of the performance of different density functionals such as B3LYP, PBE0, CAM-B3LYP and BHLYP and of *ab initio* methods such as RI-CC2 and ADC2 on the ground and excited state geometries (Tables S1 and S3 and Fig. S2 of the ESI[†]).⁴⁷ For all the DFT calculations, Grimme's dispersion corrections (D3) together with Becke and Johnson (BJ) damping were accounted for.^{48,49}

We found the best agreement with the experimental absorption spectra in toluene and DCM^{11,13} for a combination of PBE0^{50,51}–D3/def-SV(P)⁵² geometries and DFT/MRCI excitation energies and oscillator strengths. Therefore, all further calculations were performed following this computational protocol. The conductor-like screening model (COSMO) was used to mimic the solvent environment ($\epsilon = 2.380$ for toluene and $\epsilon = 8.930$ for DCM).^{53,54} The geometries of the electronic ground and excited states of **2CzPN** were all optimized without symmetry constraints. The optimized minima of the low-lying singlet excited states were obtained by using TDDFT.⁵⁵ For the triplet excited states, the Tamm–Dancoff approximation (TDA)⁵⁶ to TDDFT was employed. The TURBOMOLE program was used for all the geometry optimizations.⁵⁷ The harmonic vibrational frequency calculations were performed employing the AOFORCE module^{58,59} of TURBOMOLE for the ground state, and the SNF program⁶⁰ for the excited states.

Vertical and adiabatic excitation energies and optical electronic properties were computed using the DFT/MRCI method.³⁷ The tight parametrization of the Hamiltonian, reported in ref. 38, was employed (DFT/MRCI-R2016), which has been specially designed for multichromophoric systems. Up to 10 roots were calculated for each singlet and triplet manifold. For a better understanding of the relaxation of the system, linearly interpolated potential energy profiles (LIPs) between target singlet and triplet state minima were computed.

A wavefunction analysis of the singlet and triplet states was performed by using a Löwdin orthogonalization⁶¹ of the one-electron transition density matrix (1TDM) as implemented in the TheoDORE program of F. Plasser.⁶² This ansatz was used to characterize the nature of the electronically excited states in the singlet and triplet manifold. In this context, the triplet wavefunction analysis requires singlet–triplet transition density matrices. These densities were not available from the current DFT/MRCI code. Therefore, we extended the DFT/MRCI program to supply singlet–triplet transition density matrices and TheoDORE was modified to read these new density matrices enabling the triplet wavefunction analysis. Here, the system was decomposed into three fragments for calculating the charge transfer numbers (Ω_{AB}), charge transfer character (ω_{CT}), and natural transition orbitals (NTOs) of the Ω matrices for the individual states.^{63,64} The Cz rings of **2CzPN** constituted each one fragment (separately) and the phthalonitrile (PN) moiety the third fragment.

The spin–orbit matrix elements (SOMEs) coupling target singlet and triplet states were calculated with the spin–orbit

coupling kit (SPOCK) developed in our group.^{65–67} Herein, the spin–orbit coupling (SOC) is described by the Breit–Pauli Hamiltonian and the spin–orbit-mean-field-approximation (SOMF) is employed.^{68,69} Rate constants for ISC and RISC between singlet and triplet states were determined in the framework of the Fermi golden-rule approximation and a time-dependent approach to ISC mediated by SOC within the Condon and Herzberg–Teller approximations as implemented in the VIBES program.^{70,71} The derivatives of the SOMEs (SOMEs) with respect to mass-weighted normal coordinates were obtained as the average of two-point finite differences using a displacement of ± 0.1 along the dimensionless coordinates.⁷² For fixing the phases of the DFT/MRCI wave functions, the MO phases at each displaced geometry were set to match the ones of the equilibrium geometry. The overlap matrix between them was calculated, where the off-diagonal elements give an indication of a change of phase or ordering of the MOs. Then, the phases of the wave functions were adjusted in such a way that the largest coefficient of each wave function is positive.⁷³ Temperature effects on the rate constants were accounted for by assuming a Boltzmann distribution of the harmonic vibrational state populations in the initial electronic state.⁷⁴ Zero-point vibrational energy (ZPVE) corrections are automatically included in the VIBES calculations. For further details on the theory and the methods used for calculating ISC rate constants check the previously published review article.²⁵

3. Results and discussion

In the following we present the results obtained by our quantum chemical calculations. First, the ground state geometrical and optical properties are discussed considering also other experimental and theoretical reports. The energetics is analysed in terms of the DFT/MRCI vertical and adiabatic transition energies and interpolated pathways between target states. The characters of the low-lying excited states are analysed and the kinetics of different (R)ISC decay pathways are discussed considering direct and vibronic SOC. The Cartesian coordinates of all the computed minima, the frontier molecular orbitals (MOs) (Fig. S1 and S4–S9, ESI[†]) and electronic spectra (Tables S5–S8, ESI[†]) calculated at each of these geometries together with additional electronic structural information are provided in the ESI.[†]

3.1. Geometries and electronic structure at the S_0 state

The optimized ground state minima of **2CzPN** at the PBE0/def-SV(P) level of theory are displayed in Fig. 2a and b. Two conformers were found (S_0 and S_0'), where the difference between them is the relative orientation of the Cz groups w.r.t. the PN moiety. As expected by steric effects, a rather distorted orientation of the two bulky Cz groups is found, thus preventing the possibility of finding stable minima of higher symmetry in this emitter. The two optimized minima of **2CzPN** are not the optical reflection of one another. For the S_0' , the opening angle of the Cz rings is wider. In fact, their ground



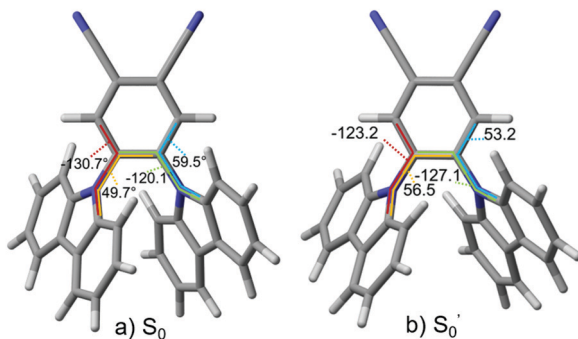


Fig. 2 Selected geometrical parameters (dihedral angles) of the ground state minima of **2CzPN** optimized at the PBE0-D3(BJ)/def-SV(P) level of theory.

state energies differ by 0.074 eV (1.7 kcal mol⁻¹), with the S_0 geometry in Fig. 2a being more stable.

The calculated geometries agree well with the experimental X-ray structures reported by M. Wong *et al.*⁷⁵ A comparison between these geometries and other S_0 minima obtained at other theoretical levels can be found in the ESI† (Table S1). We also expect enantiomers for S_0 and S_0' to exist.

The calculated DFT/MRCI-R2016 absorption spectra of both conformers of the isolated **2CzPN** are presented in Fig. 3a. The first absorption band is composed of two CT transitions. While the wavelengths of the intensive S_2 absorptions and hence the positions of the peak maxima are nearly identical in both cases, the widths of the first absorption band varies. Due to its red-shifted S_1 absorption, interaction of the S_0 conformer (Fig. 2a) with visible light yields a broader first absorption band. Also the relative intensities and energetic positions of the second absorption bands of the two conformers differ. For comparison with experimental results, the calculations were repeated in toluene and DCM environments. The DFT/MRCI-R2016 solution spectra of the S_0 conformer are shown in Fig. 3b

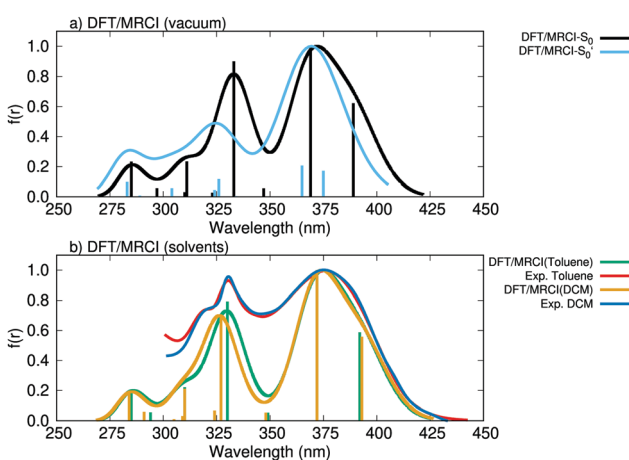


Fig. 3 DFT/MRCI-R2016 absorption spectra of **2CzPN** calculated at the S_0 minimum optimized at the PBE0-D3(BJ)/def-SV(P) level of theory: (a) in vacuum and (b) in toluene and dichloromethane (DCM) using the COSMO model. The experimental absorption spectra in toluene¹³ and DCM¹¹ are also presented.

without any empirical tuning, together with two experimental absorption spectra measured in toluene¹³ and DCM.¹¹ Solvent effects on the absorption spectra appear to be very small. Excellent agreement between the computed DFT/MRCI-R2016 spectra of the most stable S_0 conformer in Fig. 2a and the experimental data is observed. In the following, we therefore focus the analysis on this conformer.

The DFT/MRCI-R2016 vertical energies of the low-lying singlet and triplet states calculated at the S_0 state geometry of **2CzPN** are listed in Table 1. These are compared to other theoretical and experimental reports which are also displayed in this table. Experimental excitation energies and oscillator strengths (in toluene and DCM) agree well with the calculated values of the **2CzPN** in vacuum. All these excited states can be associated with $\pi\pi^*$ electronic transitions, and the corresponding frontier MOs relevant to these states are presented in Fig. 4a. From this figure, it can be observed that the H and H-2 MOs are mostly localized on the Cz units but present also some amplitudes on the PN unit. While the H-2 is mostly located on the Cz units, the H-6 contains strong localization on the PN unit and lower density in the Cz units. Regarding the MOs unoccupied in the electronic ground state, the L and L+1 MOs present amplitudes on the PN unit and at the nitrogen atom of the Cz units, while the L+2 and L+3 are mostly localized on the Cz units.

The fragmentation analysis is presented in Fig. 5b for the low-lying singlet and triplet states. This procedure will allow us to characterize the nature of the calculated DFT/MRCI-R2016 excited states of **2CzPN** in a simple manner. The lowest excited states present a considerably large CT character from the Cz substituents to the PN unit (in blue region of the bars) combined with some (lower) contributions of local excitations within the PN (in orange) unit and the Cz units (in green).

Among the low-lying singlets, there are three states with considerable oscillator strength (Table 1), namely S_1 , S_2 and S_4 . The wavefunction of the S_1 CT state of **2CzPN** is strongly dominated by an $H \rightarrow L$ electronic transition and shows up at 3.19 eV (in the gas phase), which is expected to be slightly red-shifted due to solvent-solute interactions. This value agrees well with the experiment (3.19 eV) in toluene. Here it is interesting to note that functionals like CAM-B3LYP and ω B97X-D yield excitation energies above 3.6 eV for this state.^{34,42} At 0.17 eV above, the S_2 state is found arising from a $H-2 \rightarrow L$ transition. Like S_1 , the S_2 excitation has predominantly CT character, mixed with LE on the PN unit. Together, they form the first absorption band with maximum around 375 nm. At variance with the interpretation of Hosokai *et al.*¹³ who assigned the band with maximum at approx. 325 nm to originate from $\pi\pi^*$ electronic transitions within the Cz units, our fragmentation analysis reveals the S_4 state to have mainly CT character. It is a multiconfigurational state with a leading term corresponding to a $H \rightarrow L+1$ excitation. The S_3 state, with much lower oscillator strength, corresponds to a $H-1 \rightarrow L$ transition and shows up 0.38 eV above the S_1 state.

The electronic structure of the low-lying triplet states is more mixed in the Franck-Condon (FC) region, *i.e.*, the weight

Table 1 DFT/MRCI-R2016 vertical excitation energies (ΔE_v , eV) of low-lying singlet and triplet states of the isolated **2CzPN** calculated at the S_0 minimum optimized at the PBE0-D3(BJ)/def-SV(P) level of theory

State	ΔE_v	Transition	% weight	$f(L)$	Theory ^a	Exp. ^b
S_0	0.00					
S_1	3.19	$H \rightarrow L$	78.0	0.126	2.98, ^c 3.27, ^d 3.22, ^f 3.13, ^g 3.68, ^h 3.15 ⁱ	3.19, ^e 3.30, ^j 3.32 ^k
S_2	3.36	$H-2 \rightarrow L$	80.3	0.202	3.17, ^c 3.55 ^d	
S_3	3.57	$H-1 \rightarrow L$	79.0	0.011		
S_4	3.72	$H \rightarrow L+1$	69.4	0.182		
S_5	3.83	$H-3 \rightarrow L$	81.7	0.006		
T_1	2.79	$H \rightarrow L$	59.1		2.68, ^c 3.02, ^d 2.70, ^g 2.81, ⁱ 2.50 ^m	
		$H-6 \rightarrow L$	15.1			
		$H-1 \rightarrow L$	11.7			
T_2	3.13	$H-2 \rightarrow L$	72.4		3.04 ^f	
T_3	3.24	$H-1 \rightarrow L+3$	25.9			
		$H-3 \rightarrow L+2$	24.7			
		$H-1 \rightarrow L$	14.3			
T_4	3.27	$H-1 \rightarrow L+2$	31.5			
		$H-3 \rightarrow L+3$	28.2			
		$H \rightarrow L+2$	12.5			
T_5	3.50	$H \rightarrow L+1$	28.8			
		$H \rightarrow L+2$	21.2			
		$H-2 \rightarrow L+3$	16.8			

^a Other theoretical reports. ^b Experimental reports. ^c TD-PBE0(D3)/SV(P). ^d TD-ADC(2)/SV(P). ^e Ref. 42, in cyclohexane. ^f Ref. 35, TD-LC-w*PBE/6-31+G* in toluene (PCM). ^g Ref. 42, TD-MPW1B95/6-31G*. ^h Ref. 42, TD-CAMB3LYP/6-31G*. ⁱ Ref. 45, TDA-B2(GP)-PLYP/def2-TZVP/PBE0-D3(BJ)/def2-TZVP in toluene (COSMO). ^j Ref. 13, toluene solution. ^k Ref. 16, hexane solution. ^l Ref. 46, TDA-LC-w*PBE/6-31+G* in toluene (PCM). ^m Ref. 42, TD-B3LYP/6-31G*.

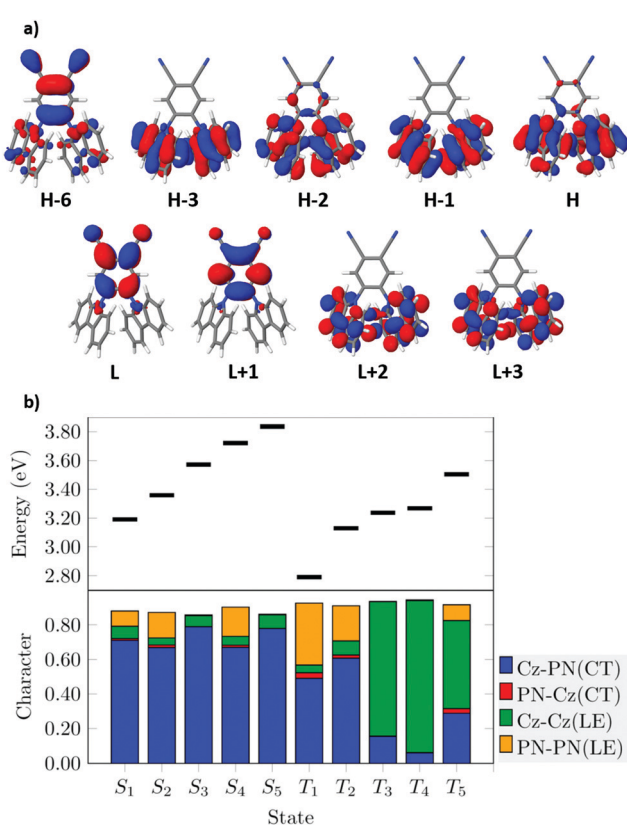


Fig. 4 (a) BH-LYP frontier molecular orbitals of **2CzPN** calculated at the S_0 geometry optimized at the PBE0-D3(BJ)/def-SV(P) level of theory. (b) Fragment-based analysis for the vertical singlet and triplet state DFT/MRCI-R2016 wavefunctions.

percentage of their dominant contributions is lower than in the corresponding singlet states and have a more combined CT –

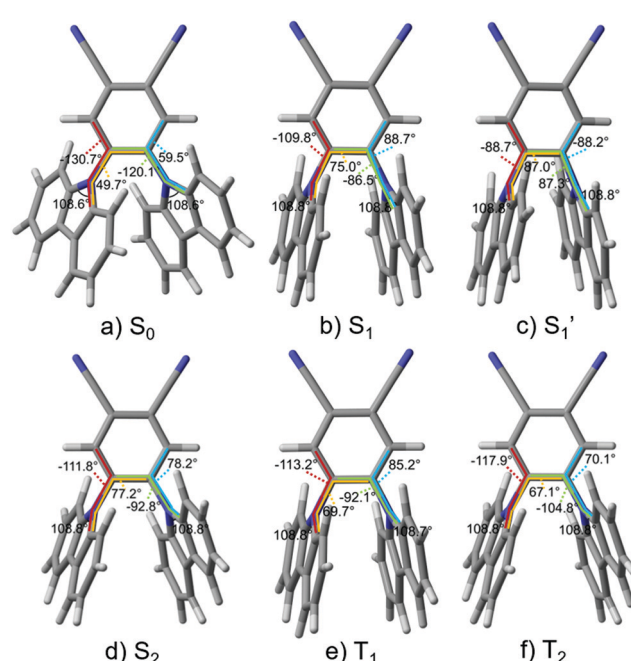


Fig. 5 Selected geometrical parameters (dihedral angles) of the ground and excited state minima of **2CzPN** optimized at the PBE0-D3(BJ)/def-SV(P) level of theory. TDDFT was used for the optimization of the excited singlet states, while TDA was used for the optimization of the triplet states.

LE character as shown by the fragmentation analysis of the transition densities (see Fig. 4b). Vertically, two triplet states below the S_1 state can be found. The T_1 state shows up at 2.79 eV ($\Delta E_{ST} = 0.40$ eV) and arises from a linear combination of three excitations: $H \rightarrow L$ (59%), $H-6 \rightarrow L$ (15%) and $H-1 \rightarrow L$ (12%). Here, the $H-6 \rightarrow L$ transition is largely occurring between MOs localized at the **PN** moiety. As such, this state



has a higher LE character compared to its singlet counterpart (S_1 state) in this geometry. Other theoretical reports have also assigned the T_1 state of **2CzPN** with a strongly mixed CT-LE character, *e.g.* by using optimally tuned long-range corrected functionals and PCM⁵⁴ and uncorrected TDA-PBE0²⁸ and TD-B3LYP³⁴ functionals. In addition, Huang *et al.*⁴² have shown how the LE contribution of this state increases with the increase of the HF exchange in conventional functionals.

Only 0.06 eV below the S_1 state is the T_2 state, characterized by a large contribution from a $H-2 \rightarrow L$ (72%) CT excitation. This energy separation is smaller than when using an empirically corrected TD-LC- ω *PBE functional and the 6-31+G* basis set (*ca.* 0.1 eV).³⁵ The following (T_3-T_5) states present a substantial LE character emerging from transitions between the **Cz** units as shown in Fig. 4.

3.2. Minima of the low-lying excited states

The energy distribution and nature of the adiabatic minima of the low-lying singlet and triplet excited states of **2CzPN** may drive the plausible energy relaxation channels, for instance, ISC and RISC. The DFT/MRCI-R2016 adiabatic excitation energies of the optimized low-lying singlet and triplet states are listed in Table 2. The corresponding geometries are presented in Fig. 5.

Minima for the S_1 , S_2 , T_1 and T_2 states were found. Two enantiomeric minima for each excited state related to the opposite orientation of the **Cz** groups in space are obtained. We have considered the two geometries for the S_1 state (S_1 and S_1') for the present analysis. In this manner, we are able to cover relaxation pathways arising from both directions of coordinate space. The minimum geometries of the S_2' , T_1' and T_2' states are given in Fig. S3 of the ESI.[†]

A common feature of these optimized excited state minima is the altered orientation of the **Cz** rings. They arrange in a more parallel fashion with respect to one another and in a more perpendicular fashion with regard to the **PN** ring. This trend is the largest for the $S_1(S_1')$ minima, followed by the T_1 and T_2 geometries (Fig. 5). Concomitantly, the CT character of these states increases upon geometry relaxation (compare Fig. 4, 6 and Fig. S10, ESI[†]). The change in electronic structure is especially pronounced for the T_1 state, which exhibits a considerable CT-LE mixing in the FC region but adiabatically it is

Table 2 DFT/MRCI-R2016 adiabatic excitation energies (ΔE^{ad} , eV) of low-lying singlet and triplet states of **2CzPN** calculated at their corresponding equilibrium geometry in vacuum

State	ΔE (eV)	Transition	% weight	ZPVE	Exp.	Theory ^b
S_1	2.91	$H \rightarrow L$	89.3	-0.09	2.94, ^a 2.96 ^e	2.84, ^c 2.99 ^d
S_1'	2.92	$H \rightarrow L$	89.4	-0.10		
S_2	3.13	$H-2 \rightarrow L$	83.1	-0.01		
T_1	2.79	$H \rightarrow L$	84.7	-0.10	2.75, ^a 2.65 ^e	2.50, ^c 2.53, ^d
T_2	3.01	$H-2 \rightarrow L$	83.4	-0.07		2.71 ^c

^a Ref. 13, onset of the fluorescence and phosphorescence (77 K) spectra in toluene. ^b Other theoretical reports. ^c Ref. 42, TD-B3LYP/6-31G*, UDFT for the triplet states. ^d Ref. 42, TD-PBE0/6-31G*, UDFT for the triplet states. ^e Ref. 7, calculated from the highest peak of the emission spectrum in toluene at 77 K.

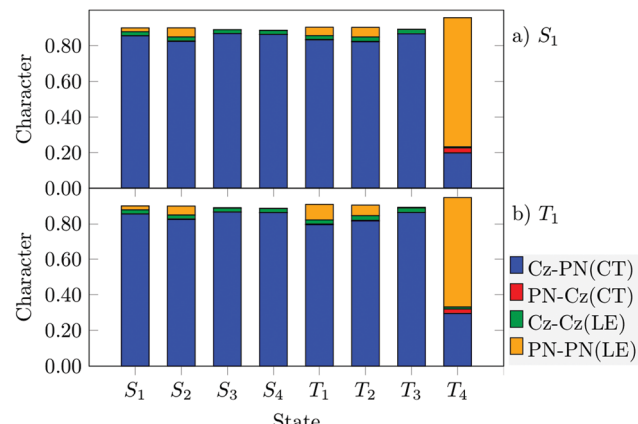


Fig. 6 Fragment-based analysis of the DFT/MRCI-R2016 wavefunctions performed at the (a) S_1 and (b) T_1 state minima.

mainly a CT state. This result is at variance with conclusions drawn by other authors who predict a much higher LE character for the adiabatic T_1 state of **2CzPN** (near 50%, LC- ω *PBE/6-31G*) on the basis of CT indices obtained by orbital composition analysis⁴² and natural transition orbital analysis³⁵.

In line with the large geometry change, the geometry relaxation effect leads to an energy stabilization of the S_1 state by *ca.* 0.28 eV w.r.t. the FC region. The onsets of the experimental emission spectra in toluene^{7,13} (2.94–2.96 eV) match the DFT/MRCI-R2016 adiabatic energy for the S_1 state of 2.91 eV very closely (Table 2). A similar stabilizing effect is found for the S_2 state (by 0.23 eV). In contrast, the geometry relaxation effect on the T_1 state energy is significantly lower than in the S_1 state, with the consequence that the ΔE_{ST} between the S_1 and T_1 adiabatic states is substantially smaller than that in the FC region. Experimental estimates of ΔE_{ST} vary between 0.09 eV and 0.35 eV, depending on the molecular environment and on the underlying model.^{5,7,10,13,76} Our adiabatic ΔE_{ST} gap of 0.12 eV (0.13 eV after inclusion of ZPVE corrections) agrees well with the experimental value in an mCP OLED (0.09 eV), estimated from peak emission fluorescence (300 K) and phosphorescence (5 K) wavelengths.¹⁰ By considering the threshold energies of the experimental fluorescence and phosphorescence bands (77 K) in toluene solution as a reference, a ΔE_{ST} gap of 0.21 eV is derived.¹³ An even larger ΔE_{ST} value of 0.31 eV is obtained if the T_1^{0-0} transition energy is taken to coincide with the first phosphorescence band maximum (77 K), while S_1^{0-0} is derived from the crossing point of the absorption and fluorescence bands at 300 K.^{7,42} Other TDDFT and TDA reports best agreeing with this (latter) value were ΔE_{ST} values of *e.g.* 0.49, 0.34 and 0.46 eV, which can be obtained when using TD- ω B97X/6-31G*,³⁰ TD-B3LYP and TD-PBE0³⁴ with the same basis set.

The T_2 CT state is influenced by a small percentage of LE character (Fig. S10, ESI[†]). Its minimum shows up 0.10 eV above the adiabatic S_1 state. This situation is different from the FC region where this state is located below the S_1 state. In contrast, TD-B3LYP calculations predict this state to be located adiabatically below the S_1 state (by 0.17 eV) being the only study reporting about the participation of this state in the (R)ISC



decay processes on **2CzPN**.³⁴ Preliminary DFT/MRCI calculations locate two adiabatic triplet states (T_1 and T_2) below the S_1 state in the **4CzIPN** TADF emitter,¹² with the T_2 state being halfway from the T_1 and S_1 states. This strongly suggests marked differences between the TADF mechanisms of both emitters: for **4CzIPN**, the presence of the intermediate T_2 state helps to accelerate the (R)ISC processes, while for **2CzPN**, pathways involving the adiabatic T_2 state are thermally activated.

3.3. Relaxation channels

To shed light on possible energy dissipation mechanisms arising upon photoexcitation, we constructed LIPs connecting the optimized electronic states as displayed in Fig. 7. These LIPs offer a qualitative view of the overall photophysics of the excited molecule. The LIPs were calculated using the respective adiabatic minimum internal coordinates.

Fig. 7a shows the LIP between the minima of the S_0 and the S_1 states. Photoexcitation at the FC point eventually results in the population of the adiabatic S_1 state due to Kasha's rule. Along this relaxation path that includes marked changes in the Cz torsional angles, we found a crossing between the S_1 and T_2 PESs in the vicinity of the FC region. A value of 0.18 cm^{-1} for their mutual SOC at this crossing point indicates that ISC is possible, in principle. However, while these two states are separated by only -0.06 eV in the FC region, their energy gap rises to 0.44 eV at the S_1 minimum. A crossing of the S_1 and T_2 PESs also occurs along a pathway connecting their adiabatic minima (Fig. 7c), in this case with a small energy barrier ($\sim 0.1 \text{ eV}$). In contrast, the T_1 and T_2

PESs do not intersect (Fig. 7f), suggesting that their vibronic coupling is small. The interconversion of the S_1 and S_1' enantiomers proceeds without a substantial barrier (Fig. 7b). As the Cz units are nearly perpendicular to the PN ring, the interconversion coordinate is mainly a sheering distortion. The S_1 and T_1 PESs show parallel tracks during this interconversion. Note the close energetic proximity of the S_1 and T_1 PESs at these geometries, a consequence of the similar nuclear arrangements at the S_1 and T_1 minima and the drastically reduced LE character of the adiabatic T_1 state in comparison to those of the FC region.

From the S_1 minimum, only the T_1 state is accessible in a downhill process, whereas the T_2 state is lying 0.10 eV above the adiabatic S_1 state. Therefore, ISC channels in **2CzPN** may arise from $S_1 \rightarrow T_1$ and $S_1 \rightarrow T_2$ decay, where the latter is thermally activated. In electroluminescent devices, RISC is expected to proceed mainly through a $T_1 \rightarrow S_1$ channel as this is the only state lying energetically close by (see Fig. 7d). However, contributions involving the T_2 state cannot be excluded *per se*. Several factors are decisive for efficient (R)ISC between target states such as the size of SOC, their adiabatic energy difference and their mutual vibrational overlap, which in turn also displays temperature dependence. In the following, the properties driving the mentioned (R)ISC channels are discussed in detail.

The component-averaged SOMEs between these target singlet and triplet states calculated at each minimum are displayed in Table 3. As expected from El-Sayed's rule, the magnitude of these SOMEs is in general small as also commonly found for transitions between $\pi\pi^*$ CT states of organic molecules. The

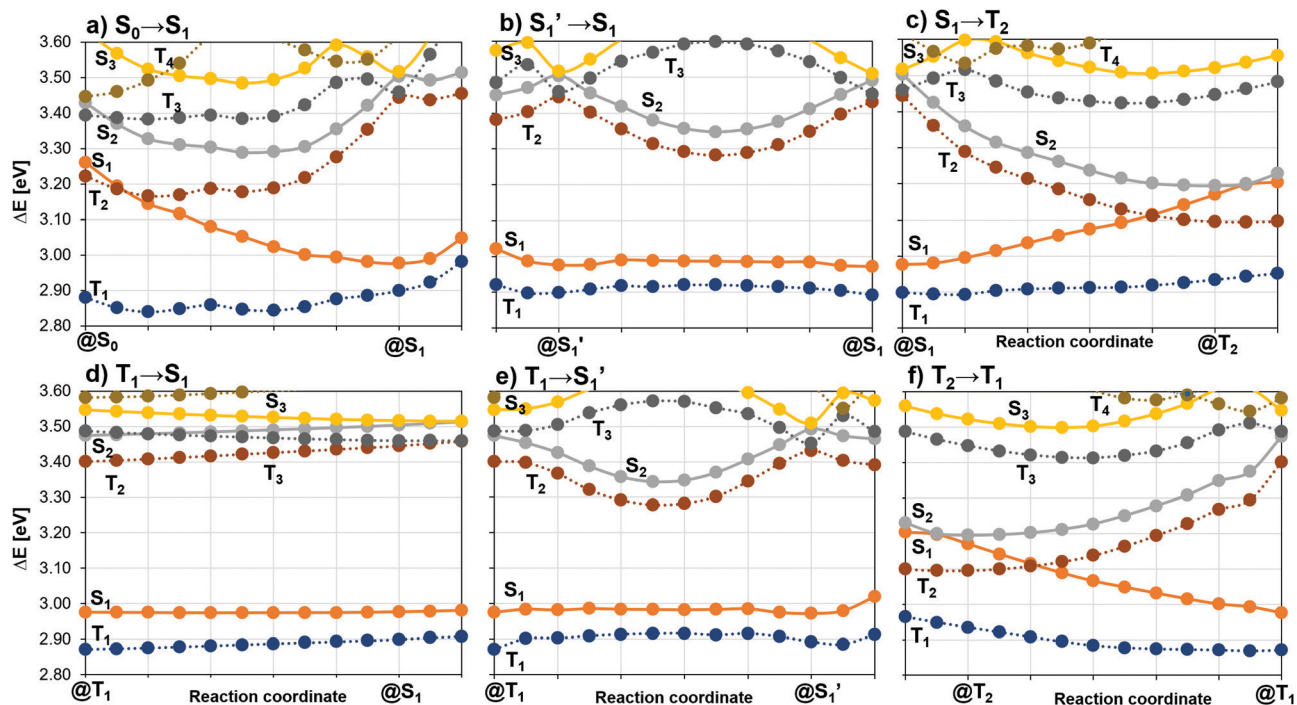


Fig. 7 DFT/MRCI-R2016 energies calculated at linear interpolated pathways (LIPs) between target electronic state minima of **2CzPN**. Dashed lines correspond to triplet state PESs, while continuous lines correspond to singlet states. Excitation energies calculated relative to the S_0 state at its corresponding minimum.



Table 3 Component-averaged spin–orbit matrix elements (SOMEs, cm^{-1}) between low-lying singlet and triplet states of **2CzPN** calculated at the optimized minima

SOMEs	@S ₀	@S ₁	@S _{1'}	@T ₁	@T ₂
$\langle T_1 \hat{H}_{\text{SO}} S_1 \rangle$	0.479	0.211	0.217	0.292	0.499
$\langle T_2 \hat{H}_{\text{SO}} S_1 \rangle$	0.183	0.223	0.249	0.240	0.228
$\langle T_3 \hat{H}_{\text{SO}} S_1 \rangle$	0.033	0.085	0.086	—	

coupling between the S_1 and T_1 CT states ($\sim 0.2 \text{ cm}^{-1}$ at their corresponding minima) arises mainly from the minor LE contributions to the triplet wavefunction (Fig. 6). The larger $\langle T_1 | \hat{H}_{\text{SO}} | S_1 \rangle$ value at the S_0 geometry correlates with the higher percentage of LE PN configurations and the increased ΔE_{ST} gap in the FC region. Other computational studies report higher SOC values at the excited-state minima as well, *e.g.*, $\text{SOME}_{S_1 \rightarrow T_1} = 0.737 \text{ cm}^{-1}$ @ T_1 minimum.³⁴ The authors mention, however, that the underlying B3LYP-TDA/6-31G* calculations significantly overestimate the S_1 and T_1 excitation energies.

The effect of vibronic SOC in **2CzPN** has been analysed in the literature by combining molecular dynamics (MD) and TDDFT-TDA calculations.^{27,28} Torsional motions modulating the CT-LE mixture were found to facilitate the SOC-driven ISC in **2CzPN**. The polarization of experimentally observed ESR signals also supported a spin-vibronic ISC mechanism in this emitter.²⁷ In contrast, a study applying time-resolved infrared vibrational spectroscopy in conjunction with (TD)DFT calculations comes to the conclusion that the suppression of structural changes upon S_1 – T_1 conversion assists the TADF process in Cz benzonitrile derivatives.¹⁵

As an alternative to non-adiabatic MD, static approaches based on Herzberg–Teller (HT)-like expansions can be used to determine spin-vibronic interactions.²⁶ To this end, derivatives of the SOMEs (∂ SOMEs) with respect to distortions along the dimensionless normal mode (v_i) displacements have to be computed. The resulting ∂ SOME values (Table S10, ESI†) are very small. For the S_1 – T_1 coupling, the maximum variation of the ∂ SOMEs is found in the second decimal place, using either the S_1 or T_1 minimum nuclear arrangements as the expansion origin, while for the S_1 – T_2 coupling a few values between 0.2 and 0.4 cm^{-1} can be found. As a consequence, vibronic contributions to the S_1 – T_1 (R)ISC rate constants are expected to be weak.

The (R)ISC rate constants ($k_{\text{ISC}}/k_{\text{RISC}}$) calculated in the framework of the FC and HT approximations at 300 K for **2CzPN** are collected in Table 4. In this table, the rate constants involving the $S_1 \rightarrow T_1$ and $T_1 \rightarrow S_1$ channels were also calculated considering the reported experimental ΔE_{ST} gaps of 0.21 eV¹³ and 0.31 eV⁷ in toluene solution. Pathways connecting different enantiomers (*e.g.* $S_1' \rightarrow T_{1,2}$ or $T_1 \rightarrow S_1'$) result in very low values for the associated (R)ISC rate constants. They are therefore not displayed in Table 4. Due to the large geometrical (spatial) difference between the initial and final states, their vibrational overlaps are very small. Instead, interconversion of the enantiomers can easily proceed on the S_1 or T_1 PESs, as shown in Fig. 7b and e.

Table 4 Computed (R)ISC rate constants $k_{(\text{R})\text{ISC}}$ (s^{-1}) for **2CzPN** (300 K) in the Condon ($k_{(\text{R})\text{ISC}}^{\text{C}}$) and Herzberg–Teller approximations ($k_{(\text{R})\text{ISC}}^{\text{HT}}$) in comparison to experimental results at room temperature

ISC	ΔE_{ST} (eV)	$k_{\text{ISC}}^{\text{C}}$ (300 K)	$k_{\text{ISC}}^{\text{HT}}$ (300 K)	k_{ISC} (exp.)
S_1 – T_1	+0.12	3.2×10^6	7.6×10^6	
	+0.09 ^a			3.1×10^7 ^a
	+0.21 ^b	8.9×10^5	3.7×10^6	2.1×10^7 ^b
	+0.29 ^c	—	—	2.9×10^7 ^c
	+0.31 ^d	5.8×10^5	1.6×10^6	3.5×10^7 ^d
	+0.35 ^e	—	—	3.0×10^7 ^e
S_1 – T_2	-0.10	1.1×10^4	9.3×10^4	
RISC	ΔE_{ST} (eV)	$k_{\text{RISC}}^{\text{C}}$ (300 K)	$k_{\text{RISC}}^{\text{HT}}$ (300 K)	k_{RISC} (exp.)
T_1 – S_1	-0.12	1.9×10^4	2.7×10^4	
	-0.09 ^a			5.6×10^3 ^a
	-0.21 ^b	1.5×10^2	4.3×10^2	6.0×10^3 ^b
	-0.29 ^c	—	—	5.4×10^3 ^c
	-0.31 ^d	2.1	2.9	1.7×10^5 ^d
	-0.35 ^e	—	—	7.9×10^4 ^e
NR ^f	—	—	—	3.5×10^6 ^f
T_2 – S_1	+0.10	9.1×10^5	—	—

^a Ref. 10, in mCP OLED. ^b Ref. 13, toluene (10^{-4} M) solution. ^c Ref. 76, co-deposited films (mCBP host matrix). ^d Ref. 7, toluene (10^{-5} M) solution. ^e Ref. 5, THF solutions. ^f Ref. 77, toluene (NR: not reported).

Vibronic SOC enhances the (R)ISC rate constants roughly by a factor of 2 at 300 K, as may be seen when comparing the $k_{(\text{R})\text{ISC}}^{\text{HT}}$ values with the direct $k_{\text{ISC}}^{\text{C}}$ values in Table 4. While an exponential decrease of the nonradiative transition rate constant may be expected from the energy gap law for weak coupling cases,²⁶ no correlation between the experimentally derived k_{ISC} and ΔE_{ST} values of **2CzPN** can be made out. All experimental studies agree that $k_{\text{ISC}} (S_1 \rightarrow T_1)$ is in the range of $2.1\text{--}3.5 \times 10^7 \text{ s}^{-1}$. In contrast, $k_{\text{RISC}} (T_1 \rightarrow S_1)$ depends substantially (varying by *ca.* three orders of magnitude) on the experimental conditions and the model for determining the kinetic constants from the measurements. The Adachi group reports $k_{\text{RISC}} (T_1 \rightarrow S_1)$ values varying slightly in the range of $5.4\text{--}6.0 \times 10^3 \text{ s}^{-1}$, despite appreciable ΔE_{ST} shifts (0.09–0.29 eV).^{10,13,76} Much higher RISC rate constants ($7.9 \times 10^4\text{--}1.7 \times 10^5 \text{ s}^{-1}$) were published by two Chinese work groups although their experimentally deduced ΔE_{ST} values (0.31–0.35 eV) are even larger.^{5,7} Our computed $S_1 \rightarrow T_1$ ISC rate constant ($7.6 \times 10^6 \text{ s}^{-1}$ at 300 K for an energy gap of $\Delta E_{\text{ST}} = 0.12 \text{ eV}$) is smaller than the experimentally determined ones by a factor of about 3 (Table 4). Unlike the experimental values, the computed ISC rate constants decrease to $1.6 \times 10^6 \text{ s}^{-1}$, if the PESs are vertically shifted to match $\Delta E_{\text{ST}} = 0.31 \text{ eV}$. The impact of the energy gap on the RISC rate constant is even more pronounced, as may be expected for TADF from the Boltzmann relation. Our computed $k_{\text{RISC}} (T_1 \rightarrow S_1)$ value of $2.7 \times 10^4 \text{ s}^{-1}$ (300 K) is approximately in the right ballpark given the high sensitivity of the results with respect to the singlet-triplet energy gap. With the increase of ΔE_{ST} from 0.12 to 0.31 eV, we observe a substantial reduction of the computed RISC rate constant by 4 orders of magnitude.

So far, we have not discussed the involvement of the T_2 state in the ISC and RISC processes. According to our calculations, T_2 is located adiabatically 0.22 eV above the T_1 state and 0.10 eV



above S_1 . Population of the T_2 state thus requires thermal activation. With a computed HT rate constant of approximately $9.3 \times 10^4 \text{ s}^{-1}$, the $S_1 \rightarrow T_2$ transition cannot compete with the much faster $S_1 \rightarrow T_1$ process. The reverse $T_2 \rightarrow S_1$ transition is a downhill process, but requires the population of the T_2 levels either by ISC from the S_1 state or by activated internal conversion from T_1 . Assuming a Boltzmann relation for the latter spin-allowed process yields a prefactor of about 10^{-4} , showing that the $T_2 \rightarrow S_1$ RISC does not essentially contribute to the TADF process unless hot excitons are available.

By using semiclassical Marcus theory to compute the Franck-Condon-weighted density of states and TDDFT-TDA potentials, Xu *et al.*³⁴ computed much higher rate constants. They obtained values of $1.3\text{--}21.1 \times 10^7 \text{ s}^{-1}$ for the $S_1 \rightarrow T_1$ ISC and of $0.2\text{--}2.8 \times 10^6 \text{ s}^{-1}$ for the $T_1 \rightarrow S_1$ RISC processes, respectively, depending on the choice of reorganization energy. The very fast $S_1 \rightarrow T_2$ transition ($k_{\text{ISC}} = 2.9 \times 10^7 \text{ s}^{-1}$) reported by these authors is caused by the fact that the T_2 state is placed adiabatically below the S_1 state in their calculations at the B3LYP-TDA/6-31G* level of theory. Interestingly, Aizawa *et al.*³⁶ have recently calculated the RISC constants for a variety of TADF emitters using a method involving the singlet-triplet crossing seam-optimized geometry (TDA-LC-BLYP/6-31+G*) and Marcus non-adiabatic theory. In agreement with our findings, the authors state that there is no crossing between the S_1 and T_1 states of **2CzPN** but that the optimized intersection corresponds to a crossing with the T_2 state. A RISC rate constant in the order of 10^3 s^{-1} together with an activation energy of *ca.* 0.25 eV has been reported for isolated **2CzPN** using their methodology.

4. Conclusions

By means of combined PBE0/SV(P) and DFT/MRCI-R2016 quantum chemical computations, we have studied in detail excited-state decay pathways and (R)ISC mechanisms of the blue TADF emitter **2CzPN**. To this end, vertical and adiabatic electronic excitation energies, vibrational frequencies and wave functions as well as (R)ISC rate constants including direct and vibronic SOC were determined.

Two minima differing by 0.074 eV (1.7 kcal mol⁻¹) in energy were located in the electronic ground state of **2CzPN**. Their nuclear arrangements differ mainly in the orientation and the opening angle of the Cz molecular planes. Based on energetic considerations and on their spectral properties, we decided to focus on the more stable conformer in the following. Enantiomers of both structures exist that are separated from their respective mirror images by high barriers in the electronic ground state.

In the FC region, two triplet excited states are located below the S_1 state with energy separations of $\Delta E(S_1\text{-}T_1) = 0.40 \text{ eV}$ and $\Delta E(S_1\text{-}T_2) = 0.06 \text{ eV}$ according to the DFT/MRCI-R2016 calculations. The excited-state transition densities, studied through fragmentation analysis of the natural transition density, revealed the nature of these states to have a strong CT component.

The admixture with LE character is substantial in the triplet manifold (*e.g.* for the T_1 state) and less pronounced in the low-lying excited singlet states. The excellent agreement of the computed absorption spectrum in vacuum with available experimental absorption spectra in toluene, THF and DCM solutions and the negligible solvent effect on the spectral properties lends confidence to the chosen computational protocol.

The calculated adiabatic DFT/MRCI-R2016 excitation energies in vacuum locate only the T_1 state below the S_1 state (by 0.12 eV), while the T_2 state lies 0.10 eV above. The intersection of the S_1 and T_2 PESs occurs close to the FC point. Considering the small magnitude of the electronic SOC (component averaged $\langle T_2 | \hat{H}_{\text{SO}} | S_1 \rangle = 0.18 \text{ cm}^{-1}$) renders the dynamic ISC $S_1 \rightarrow T_2$ along the S_1 geometry relaxation path not very likely. At the optimized equilibrium structures of the S_1 and T_1 states, the Cz rings are markedly more perpendicular to the PN plane than in the ground state. The electronic decoupling of their π -systems facilitates the charge separation in these donor-acceptor systems, thus enhancing their CT character. Moreover, it enables a nearly barrierless interconversion of the enantiomers in the first excited singlet and triplet states by a sheering motion of the Cz units. Because of the small overlaps of the vibrational wavefunctions of the two enantiomeric potential energy wells, we nevertheless have indications that cross-couplings, such as $S_1' \rightarrow T_1$ ISC, do not play an important role among the excited-state decay processes.

The mutual (direct) SOCs between the S_1 and T_1 states at their respective minimum geometries are reduced in comparison to the FC point, a direct consequence of the varying LE contributions to the wave functions. Furthermore, the calculated derivatives of the $S_1\text{-}T_1$ SOMEs with respect to dimensionless normal mode displacements are small, but not negligible indicating that the contributions of Herzberg-Teller vibronic SOC to the $k_{(\text{R})\text{ISC}}$ constants are moderate. Spin-vibronic interactions are found to speed up the $S_1\text{-}T_1$ ISC and $T_1\text{-}S_1$ RISC processes by a factor of about 2. The triplet state formation is predicted to proceed predominantly *via* the $S_1 \rightarrow T_1$ channel with a rate constant of $8 \times 10^6 \text{ s}^{-1}$ at room temperature, somewhat smaller than the experimentally derived values of $2\text{--}3 \times 10^7 \text{ s}^{-1}$,^{5,7,10,13,76} but in the right ballpark. In addition, a thermally activated $S_1 \rightarrow T_2$ channel is available, but predicted to be less efficient ($k_{\text{ISC}} = 9 \times 10^4 \text{ s}^{-1}$).

Our calculated RISC rate constant associated with the back population of the S_1 state of **2CzPN** in vacuum through the $T_1\text{-}S_1$ channel ($k_{\text{RISC}} = 3 \times 10^4 \text{ s}^{-1}$) certainly is one order of magnitude larger than the smallest constants ($5\text{--}6 \times 10^3 \text{ s}^{-1}$) reported in solution and in OLEDs,^{10,13,76} but significantly smaller than the highest value ($3.5 \times 10^6 \text{ s}^{-1}$)⁷⁷ determined by kinetic modelling of transient triplet absorption spectra in toluene solution. Astonishingly, the experimentally derived RISC rate constants and ΔE_{ST} values do not appear to be correlated. On one hand, nearly constant k_{RISC} values have been reported, despite appreciably different ΔE_{ST} gaps (0.09–0.29 eV).^{10,13,76} On the other hand, a variation of experimentally derived rate constants by *ca.* 2 orders of magnitude can be made out when comparing different measurements of



2CzPN RISC in toluene solutions with similar singlet-triplet energy gaps of 0.31 and 0.29 eV, respectively.^{7,13} Following this consideration, we repeated the (R)ISC rate constant calculations using ΔE_{ST} values of two of these experimental reports, *i.e.* 0.21¹³ and 0.31 eV^{7,42} in addition to our computed value of 0.12 eV to test the impact of singlet-triplet separation on the transition probabilities. As expected for a thermally activated process in the weak coupling limit, the RISC rate constant decreases exponentially (by *ca.* 4 orders of magnitude) when increasing ΔE_{ST} from 0.12 eV to 0.31 eV, while keeping the electronic coupling terms constant. The downhill $S_1 \rightarrow T_1$ ISC process is less affected (less than one order of magnitude) by this variation of the energy gap.

Although the T_2 state lies adiabatically close to S_1 , it is not expected to play a major role in mediating the spin-flips in **2CzPN** unless the T_2 state is directly populated by hot excitons from the embedded OLED layer. Starting from the equilibrated S_1 state, the $S_1 \rightarrow T_2$ ISC is an activated process that cannot compete with the much faster $S_1 \rightarrow T_1$ ISC. Moreover, we found no indication for a strong vibronic coupling of the T_2 and T_1 potentials that could help overcome the negative exponential dependence of the RISC rate constant on the magnitude of the energy gap.

In comparison, DFT/MRCI calculations on the very efficient **4CzIPN** TADF emitter¹² show a significant CT-LE mixing in the low-lying triplet states and minor effects in the S_1 state. This complements with a T_2 state energetically located halfway from the S_1 and T_1 state, acting as an intermediate between these states, thereby accelerating the (R)ISC processes.

Our calculations show that the (R)ISC probability in **2CzPN** depends mainly on the interplay between the S_1 and T_1 states with spin-vibronic interactions accelerating the nonradiative transitions only marginally. One possible explanation for the strange findings with regard to the RISC rate constants in relation to the S_1-T_1 energy gap in **2CzPN** is that the models employed for deriving ΔE_{ST} values from spectroscopic data are not adequate. Could, alternatively, a larger singlet-triplet splitting, caused by higher LE contributions to the excited-state wave functions, be compensated by an increase of the electronic SOC strength? In principle, this is possible but difficult to imagine in this particular case. At the S_0 geometry, where the $\Delta E(S_1-T_1)$ is relatively large (0.40 eV), the magnitude of $\langle T_1 | \hat{H}_{SO} | S_1 \rangle$ differs by a factor of *ca.* 2 from the values at the relaxed excited-state geometries. The increased electronic coupling would lead to an acceleration of the ISC probability by a factor of about 4 in FC approximation, as opposed to a deceleration by 4 orders of magnitude caused by the reduced vibrational overlaps of the shifted potentials. The most plausible explanation for the high RISC rates, reported in some experimental studies,^{5,7,77} is the participation of other processes such as triplet-triplet annihilation or hot exciton transfer in the delayed fluorescence.

Conflicts of interest

There are no conflicts to declare.

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